

- 1 -

## METHOD AND KIT FOR THE CHARACTERIZATION OF ANTIBIOTIC-RESISTANCE MUTATIONS IN MYCOBACTERIUM TUBERCULOSIS

#### **DESCRIPTION**

#### Field of the Invention

This application relates to a method and kit for the characterization of antibiotic resistance mutations in *Mycobacterium tuberculosis*, and particularly to the evaluation of such mutations in clinical samples.

#### Background of the Invention

M. tuberculosis can be resistant to all antibiotics that are currently used to treat tuberculosis patients. Antibiotic resistance is due to acquired point mutations in target genes in the genome of M. tuberculosis. These point mutations render the organism insensitive to the action of the antibiotic by preventing it's uptake or activation, or by altering the antibiotic target. The observed antibiotic resistance in M. tuberculosis is not due to an episome-encoded resistance gene transferred from one strain to another and, like other bacteria, is single-step (one point mutation), high-level resistance.

Rapid and accurate detection of antibiotic resistance in *Mycobacterium* tuberculosis in sputum samples would greatly improve both patient treatment and outcome. Presently, analysis of *M. tuberculosis* is carried out on DNA recovered from sputum samples handled according to Standard Infectious Disease/Public Health Laboratory practices. The sputum sample is decontaminated and a cell sediment isolated. This cell sediment is the sample source for all routine procedures used in the detection and isolation of *M. tuberculosis*. Portions of this sample are used in BacTec cultures for selective growth of *M. tuberculosis*, agar plate/agar slant cultures for *M. tuberculosis*, acid-fast bacilli (AFB) smears for mycobacteria detection and molecular biological methods for the detection of M. tuberculosis and atypical mycobacteria. (See Fig. 1)

Mycobacterial DNA is prepared directly from the decontaminated sputum cell sediments according to standard procedures and this mycobacterial DNA is used in the various molecular biological detection procedures. The methods presently in use for the detection of *M. tuberculosis* are either PCR-based or probe-based. These tests are used

primarily on AFB smear-positive samples. Since the presence of M. Inherculosis has already been established by the AFB smear, these tests are used primarily in a confirmatory capacity as opposed to a diagnostic capacity. Furthermore, these tests provide no information on the potential antibiotic resistance of these M. inherculosis samples.

Below is a list of ambiotics used to treat M. Tuherculosts infections. The gene target of the specific antibiotic and regions associated with antibiotic resistance are listed, if known. The references on which the codon assignments are based are listed at the end of the application.

	1.	Rifampin	rpoB gene	codon 507-533°
	2.	Isoniazid	katG gene	codon 275/315/328 <sup>b</sup>
J	3.	Teoniazid	mabA gene	unknown a
	4.	Isoniazid	oxyR-ahpC intergenic region	(PR)
C) CN				nucleotides -48 to +33
ļ.	<b>5</b> .	Azithroniycin	236 rRNA sequence	nucleatide 2568A.
O.	6.	Pyrazinamide	pncA gene	codon 47/85 <sup>r</sup>
≅ .	7.	Ethambutol	embB gene	codan 306 g
O	8.	Streptomycin	rpsL/s12 gene	cadon 43/88 h
ĨIJ	9.	Streptomycin	16S/rrs sequence	nucleotides 491, 512, 516, 513,
(A Ca				903, 904 .
ļ.	10.	Ciprofloxacin	gyrA gene	codon 88-95 <sup>j</sup>

Probe-based tests do exist for the determination of tifampin resistance in M tuberculasis (line probe assay-InnoTek), but these probes rely on prior knowledge of antibiotic resistance-associated mutations in the rpoB gene. Mutations outside the region covered by the probe or new mutations not included in the probe cocktail could still confer resistance, but would not be detected using this product in it's present form.

Abstracts of the Interscience Conference on Antimicrobial Agents and Chemotherapy", vol. 34, page 163 (1994) describes the application of automated DNA sequence analysis of hap65 to speciation of isolates previously-identified as being M. tuberculosis.

Thus, there remains a need for a method for detecting antibiotic-resistance mutations in clinical *M. tuberculosis* sputum samples which is capable of detecting mutations in all of the gene targets which confer antibiotic resistance. It is an object of the present invention to provide such a method. It is a further object of this invention to provide amplification and cycle sequencing primer sets, and kits containing such primer sets, for use in the characterization of antibiotic resistance mutations in *M. tuberculosis*.

### Summary of Invention

Amplification and cycle sequencing primer sets have been developed for the detection and analysis of antibiotic resistance-associated mutations in defined regions of the rpoB (rifampin), katG (isoniazid), oxyR-ahpC PR (isoniazid), mabA (isoniazid), rpsL/s12 (streptomycin), 16S/rrs (streptomycin), embB (ethambutol), pncA (pyrazinamide), gyrA (ciprofloxacin) and 23S (azithromycin) genes. Using these primer sets and the OPENGENETM automated DNA sequencing system, a protocol has been developed which permits both the rapid detection of M. tuberculosis and the identification of antibiotic resistance-associated mutations in a series of gene targets. The present invention uses a series of tests designed to detect antibiotic resistance-associated mutation in all gene targets for all antibiotics presently in use for the treatment of tuberculosis. The tests are employed in a hierarchical manner on both AFB smear-positive or smear-negative samples to determine both the presence and antibiotic-resistance of M. tuberculosis in a given sample. This method permits the simultaneous determination of M. tuberculosis presence in a sample and the antibiotic-resistance profile to an entire panel of antibiotics. Standard methods require from 2-6 weeks to culture M. tuberculosis and additional time to establish antibiotic resistance. Although DNA sequence-based (genotypic) tests are not intended to replace the traditional culture-based (phenotypic) methods, these tests do represent a rapid, sensitive and accurate protocol which provides clinicians with valuable information regarding antibiotic treatment options within days as opposed to weeks.

### Brief Description to the Figures

Fig. 1 shows known testing protocols for M. tuberculosis; and

- 4 -

Fig. 2 shows a hierarchical assay scheme for evaluating *M. tuberculosis* type in accordance with the invention.

### **Detailed Description of the Invention**

In accordance with the invention, regions of the genome of *M. tuberculosis* associated with antibiotic resistance are amplified and sequenced using specifically designed amplification and sequencing primers. Various techniques for amplification are known, including the basic PCR amplification techniques described in US Patent No. 4,683,202, which is incorporated herein by reference. Similarly, various techniques for sequencing are know, some of which require prior amplification and some of which do not. Included among known sequencing techniques are those disclosed in US Patents Nos. 5,834,189 and 5,789,168, which are incorporated herein by reference. The primers of the invention can be used in any of these sequencing formats, although the invention is exemplified below using separate amplification and cycle-sequencing steps.

In theory, the selection of primers to amplify and sequence a known region of interest should be straightforward. In fact, however, because of the possibility of primer binding to other sites, complications arising from secondary structure, and other factors which are not fully understood, some primers perform better than others for amplification and sequencing of the same region of interest. The present invention provides primers which have been optimized for the amplification and sequencing of regions associated with each of the ten known types of antibiotic resistance. These primer sets are shown below, along with the sequence of the genes that they are used to analyze. In the gene sequences, the locations of the primers are underlined.

#### **Primers**

#### rpoB (rifampin resistance)

rpoB-F amplification primer, 20-mer, bp2201-2220

5' TAC GGT CGG CGA GCT GAT CC 3'

SEQ. ID NO. 1

rpoB-R amplification primer, 20-mer, bp2611-2592

5' TAC GGC GTT TCG ATG AAC CC 3'

- 5 -

rpoB-5s sequencing primer, 20-mer, bp2201-2220
5' TAC GGT CGG CGA GCT GAT CC 3'

rpoB-3s sequencing primer, 20-mer, bp2611-2592
5' TAC GGC GTT TCG ATG AAC CC 3'

SEQ ID NO. 4

### SEQ. ID. NO. 5

2161	aaaccgacga	catcgaccac	ttcggcaacc	gccgcctgcg	<u>tacggtcggc</u>	gagctgatcc
2221	aaaaccagat	ccgggtcggc	atgtcgcgga	tggagcgggt	ggtccgggag	cggatgacca
2281	cccaggacgt	ggaggcgatc	acaccgcaga	cgttgatcaa	catccggccg	gtggtcgccg
2341	cgatcaagga	gttcttcggc	accagccagc	tgagccaatt	catggaccag	aacaacccgc
2401	tgtcggggtt	gacccacaag	cgccgactgt	cggcgctggg	gcccggcggt	ctgtcacgtg
2461	agcgtgccgg	gctggaggtc	cgcgacgtgc	acccgtcgca	ctacggccgg	atgtgcccga
2521	tcgaaacccc	tgaggggccc	aacatcggtc	tgatcggctc	gctgtcggtg	tacgcgcggg
2581	tcaacccgtt	cgggttcatc	gaaacgccgt	accgcaaggt	ggtcgacggc	gtggttagcq

### katG (isoniazid resistance)

katG-F amplification primer, 20-mer, bp722-741

5' ATG GGG CTG ATC TAC GTG AA 3' SEQ ID NO. 6

katG-R amplification primer, 20-mer, bp1250-1231

5' GGT GTT CCA GCC AGC GAC GC 3' SEQ ID NO. 7

katG-5s sequencing primer, 20-mer, bp722-741

5' ATG GGG CTG ATC TAC GTG AA 3' SEQ ID NO. 8

katG-3s sequencing primer, 20-mer, bp1250-1231

5' GGT GTT CCA GCC AGC GAC GC 3' SEQ ID NO. 9

661	gctcggcgat	gagcgttaca	gcggtaagcg	ggatctggag	aacccgctgg	ccgcggtgca
721	gatggggctg	atctacgtga	acccggaggg	gccgaacggc	aacccggacc	ccatggccgc
781	ggcggtcgac	attcgcgaga	cgtttcggcg	catggccatg	aacgacgtcg	aaacagcggc
841	gctgatcgtc	ggcggtcaca	ctttcggtaa	g <b>acc</b> catggc	gccggcccgg	ccgatctggt
901	cggccccgaa	cccgaggctg	ctccgctgga	gcagatgggc	ttgggctgga	agagctcgta
961	tggcaccgga	accggtaagg	acgcgatcac	c <b>agc</b> ggcatc	gaggtcgtat	ggacgaacac
1021	cccgacgaaa	<b>tgg</b> gacaaca	gtttcctcga	gatcctgtac	ggctacgagt	gggagctgac

1081 gaagagccct gctggcgctt ggcaatacac cgccaaggac ggcgccggtg ccggcaccat 1141 cccggacccg ttcggcgggc cagggcgctc cccgacgatg ctggccactg acctctcgct 1201 gcgggtggat ccgatctatg agcggatcac gcgtcgctgg ctggaacacc ccgaggaatt 1261 ggccgacgag ttcgccaagg cctggtacaa gctgatccac cgagacatgg gtcccgttgc

- 6 -

### oxyR-aphC intergenic region (PR)

PR-F amplification primer, 20-mer, bp451-470

5' ACC ACT GCT TTG CCG CCA CC 3' SEQ ID NO. 11 PR-R amplification primer, 20-mer, bp687-668 5' CCG ATG AGA GCG GTG AGC TG 3' SEQ ID NO. 12 PR-5s sequencing primer, 20-mer, bp451-470 5' ACC ACT GCT TTG CCG CCA CC 3' SEQ ID NO. 13 PR-3s sequencing primer, 20-mer, bp687-668 5' CCG ATG AGA GCG GTG AGC TG 3' SEQ ID NO. 14

#### SEQ ID NO. 15

361 atgccctggg ggtgcaccga gaccggcttc cgaccaccgc tcgccgcaac gtcgactggc 421 tcatatcgag aatgcttgcg gcactgctga accactgctt tgccgccacc gcggcgaacg 481 cgcgaagccc ggccacggcc ggctagcacc tcttggcggc gatgccgata aatatggtgt 541 gatatatcac ctttgcctga cagcgacttc acggcacgat ggaatgtcgc aaccaaatgc 601 attgtccgct ttgatgatga ggagagtcat gccactgcta accattggcg atcaattccc 661 cgcctaccag ctcaccgctc tcatcggcgg tgacctgtcc aaggtcgacg ccaagcagcc 721 cggcgactac ttcaccacta tcaccagtga cgaacaccca ggcaagtggc gggtggtgtt

## mabA (isoniazid resistance)

mabA-F amplification primer, 20-mer, bp56-75	
5' CCT CGC TGC CCA GAA AGG GA 3'	SEQ ID NO. 16
mabA-R amplification primer, 20-mer, bp303-284	
5' ATC CCC CGG TTT CCT CCG GT 3'	SEQ ID NO. 17
mabA-5s sequencing primer, 20-mer, bp56-75	
5' CCT CGC TGC CCA GAA AGG GA 3'	SEQ ID NO. 18

- 7 -

mabA-3s sequencing primer, 20-mer, bp303-284

### 5' ATC CCC CGG TTT CCT CCG GT 3'

SEQ ID NO. 19

### SEQ ID NO. 20

1	agcgcgacat	acctgctgcg	caattcgtag	ggcgtcaata	cacccgcagc	caggg <u>cct.cg</u>
61	ctgcccagaa	<u>aggga</u> tccgt	catggtcgaa	gtgtgctgag	tcacaccgac	aaacgtcacg
121	agcgtaaccc	cagtgcgaaa	gttcccgccg	gaaatcgcag	ccacgttacg	ctcgtggaca
181	taccgatttc	ggcccggccg	cggcgagacg	ataggttgtc	ggggtgactg	ccacagccac
241	tgaaggggcc	aaacccccat	tcgtatcccg	ttcagtcctg	gttaccggag	gaaaccgggg
301	<u>gat</u> cgggctg	gcgatcgcac	agcggctggc	tgccgacggc	cacaaggtgg	ccgtcaccca

## rpsL/s12 (streptomycin resistance)

s12-F amplification	primer,	20-mer,	bp1-20
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5' CGG TAG ATG CCA ACC ATC CA 3' SEQ ID NO. 21

s12-R amplification primer, 20-mer, bp384-365

5' GCA TCA GCC CTT CTC CTT CT 3' SEQ ID NO 22

s12-5s sequencing primer, 20-mer, bp1-20

5' CGG TAG ATG CCA ACC ATC CA 3' SEQ ID NO. 23

s12-3s sequencing primer, 20-mer, bp384-365

5' GCA TCA GCC CTT CTC CTT CT 3' SEQ ID NO. 24

### SEQ ID NO. 25

1	cagtagatac	caaccatcca	gcagctggtc	cgcaagggtc	gtcgggacaa	gatcagtaag
61	gtcaagaccg	cggctctgaa	gggcagcccg	cagcgtcgtg	gtgtatgcac	ccgcgtgtac
121	accaccactc	cgaagaagcc	gaactcggcg	cttcggaagg	ttgcccgcgt	gaagttgacg
181	agtcaggtcg	aggtcacggc	gtacattccc	ggcgagggcc	acaacctgca	ggagcactcg
241	atggtgctgg	tgcgcggcgg	ccgggtgaag	gacctgcctg	gtgtgcgcta	caagatcatc
301	cgcggttcgc	tggatacgca	gggtgtcaag	aaccgcaaac	aggcacgcag	ccgttacggc
361	gctaagaagg	agaagggctg	atgccacqca	aggggcccgc	gcccaagcgt	ccattaatca

### 16S/rrs (streptomycin resistance)

16S-F amplification primer, 21-mer, bp5-25

5' GGT GAT CTG CCC TGC ACT TCG 3'

SEQ ID NO. 26

16S-R amplification primer, 21-mer, bp147-127

WO 00/36142

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5' CGT CAC CCC ACC AAC AAG CTG 3'	SEQ ID NO. 27
16S-5s sequencing primer, 21-mer, bp5-25	
5' GGT GAT CTG CCC TGC ACT TCG 3'	SEQ ID NO. 28
16S-3s sequencing primer, 21-mer, bp147-127	
5' CGT CAC CCC ACC AAC AAG CTG 3'	SEQ ID NO. 29

# SEQ ID NO. 30

- cgtgggtgat ctgccctgca cttcgggata agcctgggaa actgggtcta ataccggata
- 61 ggaccacggg atgcatgtct tgtggtggaa agcgctttag cggtgtggga tgagcccgcg
- 121 gcctatcagc ttgttggtgg ggtgacg

## embB (ethambutol resistance)

embB-F amplification primer, 21-mer, bp7/61-7781	
5' CGG CAA GCT GGC GCA CCT TCA 3'	SEQ ID NO. 31
embB-R amplification primer, 21-mer, bp8040-8020	
5' AGC CAG CAC ACT AGC CCG GCG 3	SEQ ID NO. 32
embB-5s sequencing primer, 21-mer, bp7761-7781	
5' CGG CAA GCT GGC GCA CCT TCA 3'	SEQ ID NO. 33
embB-3s sequencing primer, 21-mer, bp8040-8020	
5' AGC CAG CAC ACT AGC CCG GCG 3	SEO ID NO. 34

# SEQ ID NO. 35

7741	cggcatgcgc	cggctgattc	cggcaagctg	gcgcaccttc	accctgaccg	acgccgtggt
7801	gatattcggc	ttcctgctct	ggcatgtcat	cggcgcgaat	tcgtcggacg	acggctacat
7861	cctgggcatg	gcccgagtcg	ccgaccacgc	cggctacatg	tccaactatt	tccgctggtt
7921	cggcagcccg	gaggatccct	tcggctggta	ttacaacctg	ctggcgctga	tgacccatgt
7981	cagcgacgcc	agtctgtgga	tgcgcctgcc	agacctggc <u>c</u>	gccgggctag	tatactaact

# pncA (pyrazinamide resistance)

pncA-F amplification primer, 20-mer, bp1-20 5' ATG CGG GCG TTG ATC ATC GT 3'

WO 00/36142 PCT/CA99/01177 . . .

-9-

pncA-F amplification primer, 20-mer, bp561-542	
5' TCA GGA GCT GCA AAC CAA CT 3'	SEQ ID NO. 37
pncA-5s sequencing primer, 20-mer, bp1-20	
5' ATG CGG GCG TTG ATC ATC GT 3'	SEQ ID NO. 38
pncA-3s sequencing primer, 20-mer, bp561-542	
5' TCA GGA GCT GCA AAC CAA CT 3'	SEQ ID NO. 39

# SEQ ID. NO. 40

1	atgcgggcgt	tgatcatcgt	cgacgtgcag	aacgacttct	gcgagggtgg	ctcgctggcg
61	gtaaccggtg	gcgccgcgct	ggcccgcgcc	atcagcgact	acctggccga	agcggcggac
121	taccatcacg	tcgtggcaac	caaggacttc	cacatcgacc	cgggtgacca	cttctccggc
181	acaccggact	attcctcgtc	gtggccaccg	cattgcgtca	gcggtactcc	cggcgcggac
241	ttccatccca	gtctggacac	gtcggcaatc	gaggcggtgt	tctacaaggg	tgcctacacc
301	ggagcgtaca	gcggcttcga	aggagtcgac	gagaacggca	cgccactgct	gaattggctg
361	cggcaacgcg	gcgtcgatga	ggtcgatgtg	gtcggtattg	ccaccgatca	ttgtgtgcgc
421	cagacggccg	aggacgcggt	acgcaatggc	ttggccacca	gggtgctggt	ggacctgaca
481	gcgggtgtgt	cggccgatac	caccgtcgcc	gcgctggagg	agatgcgcac	cgccagcgtc
541	gagttggttt	gcagctcctg	a			

# gyrA (fluoroquinilone/ciprofloxacin resistance)

gyrA-F amplification primer, 20-mer, bp2383-2402	
5' CAG CTA CAT CGA CTA TGC GA 3'	SEQ ID NO. 41
gyrA-R amplification primer, 20-mer, bp2702-2683	
5' GGG CTT CGG TGT ACC TCA TC 3'	SEQ ID NO. 42
gyrA-5s sequencing primer, 20-mer, bp2383-2402	
5' CAG CTA CAT CGA CTA TGC GA 3'	SEQ ID NO. 43
gyrA-3s sequencing primer, 20-mer, bp2702-2683	
5' GGG CTT CGG TGT ACC TCA TC 3'	SEQ ID NO. 44

2341	cgaccggatc	gaaccggttg	acatcgagca	ggagatgcag	cgcagctaca	tcgactatgc
2401	gatgagcgtg	atcgtcggcc	gcgcgctgcc	ggaggtgcgc	gacgggctca	agcccgtgca
2461	tcgccgggtg	ctctatgcaa	tgttcgattc	cggcttccgc	ccggaccgca	gccacgccaa

2521 gtcggcccgg tcggttgccg agaccatggg caactaccac ccgcacggcg acgcgtcgat 2581 ctacgacagc ctggtgcgca tggcccagcc ctggtcgctg cgctacccgc tggtggacgg 2641 ccagggcaac ttcggctcgc caggcaatga cccaccggcg gcgatgaggt acaccgaagc 2701 ccggctgacc ccgttggcga tggagatgct gagggaaatc gacgaggaga cagtcgattt

### 23S (macrolide/azithromycin resistance)

23S-F amplification primer, 20-mer, bp2444-2463

5' CGA AAT TCC TTG TCG GGT AA 3'	SEQ ID NO. 46
23S-R amplification primer, 20-mer, bp2683-2664	
5' GTA TTT CAA CAA CGA CTC CA 3'	SEQ ID NO. 47
23S-5s sequencing primer, 20-mer, bp2444-2463	
5' CGA AAT TCC TTG TCG GGT AA 3'	SEQ ID NO. 48
23S-3s sequencing primer, 20-mer, bp2683-2664	
5' GTA TTT CAA CAA CGA CTC CA 3'	SEQ ID NO. 49

#### SEQ ID NO. 50

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2401 gccccagtaa acggcggtgg taactataac catcctaagg tagcgaaatt ccttgtcggg
2461 taagttccga cctgcacgaa tggcgtaacg acttcccaac tgtctcaacc atagactcgg
2521 cgaaattgca ctacgagtaa agatgctcgt tacgcgcggc aggacgaaaa gaccccggga
2581 ccttcactac aacttggtat tggtgttcgg tacggtttgt gtaggatagg tgggagactt
2641 tgaagcacag acgccagttt gtgtggagtc gttgttgaaa taccactctg atcgtattgg
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To facilitate detection of the sequencing products using real-time fluorescence-based electrophoresis apparatus (for example, a Visible Genetics OPENGENE<sup>TM</sup> sequencer), at least one of the sequencing primers is preferably labeled with a flourescent label. The label is selected for compatibility with the sequencing apparatus employed, and may be, for example, fluorescein or a cyanine dye such as CY5.0 OR CY5.5.

The primers of the invention are suitably packaged in a kit. This kit will contain individually packaged amplification and sequencing primers sets for each resistance gene to be evaluated by the kit. Thus, the kit of the invention includes at least 4 primers (two amplification and two sequencing primers), and preferably includes the primer sets for a plurality of resistance genes, most preferably the primer sets for all ten resistance genes.

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- 11 -

The suitable protocol for the utilization of these primer sets in the evaluation of *M. tuberculosis* in clinical samples utilizes PCR amplification, followed by cycle sequencing. DNA for use in the test is obtained from a sample of sputum (100ul-10ml). The sputum sample is processed according to Standard Infectious Disease/Public Health Laboratory practices (Mycobacteriology Bench Manual, Laboratory Services Branch, December 1997, Ontario Ministry of Health). The sputum sample is homogenized, decontaminated and concentrated. Mycobacterial DNA is prepared directly from a portion of the concentrated cell sediment (100-200ul) using standard DNA extraction methods or commercially available kits.

Amplification of the DNA is performed using the amplification primer sets described above. PCR reagents can be prepared for individual reactions, or may be prepared as a master mix which can be used for multiple tests e.g., 10 PCR reactions. Exemplary combinations of reagents are summarized in the following table.

PCR mix		1 PCR		10 PCRs	final conc. / PCR
genomic DNA	(20ng/ul)		1.0ul		20ng
(~0.5fM)					
10X PCR buffer I		2.5ul		25.0ul	1X
2.5mM dNTP mix	(1:1:1:1)	2.5ul		25.0ul	250uM
DMSO			1.3ul	13.0ul	5%
Taq DNA polymerase (1U)		0.2ul		2.0ul	1 unit
molecular grade water		16.5ul		165.0ul	
MTB gene primers	(10uM)	1.0ul	·	10.0ul	10pmol per primer
total volume per PCR	25.0ul				

If the master mix as shown in the column labeled 10 PCRs is utilized, the mastermix contains all the necessary PCR reagents other than the genomic DNA. In this example, 24.0ul of the mastermix is added to a PCR tube, that already contains 1.0ul of genomic DNA, prior to the addition of the mineral oil overlay and placement in the thermocycler.

The genomic DNA preparation utilized must be of sufficient quality and integrity for robust and reproducible PCR. Suitable DNA preparation can be obtained using

the Gentra Puregene™ DNA isolation kit. The kit components are appropriate for the isolation of genomic DNA from blood, fresh or frozen tissue, archival material and paraffinembedded tissue.

Each primer pair is used to amplify a single gene region under the following conditions:

1.	Denaturation	94°C	5 minutes	1 cycle
2.	Denaturation	94°C	30 seconds	
	Annealing	60°C	30 seconds	35 cycles
	Extension	72°C	60 seconds	
3.	Extension	72°C	5 minutes	1 cycle
4.	Hold	6°C		

The temperature change during the cycles of the step 2 is desirably set to ramp at a rate of 1°C/sec.

After amplification, 2.0ul from the 25.0ul PCR is analysed for purity on a 0.8% agarose gel. Samples displaying single PCR product bands can be used directly for sequence analysis. The yield and purity of the PCR product determines the amount to be used in the subsequent cycle sequencing reaction. Comparable verification of sequencing purity is performed on each of the other amplification products.

Sequence analysis is carried out on the amplified product. The basic procedures and conditions are the same for each region. Accordingly, the invention will be exemplified using the rpoB gene.

For initial sequence analysis of rpoB, the rpoB-5s primer should be used. For confirmatory sequence analysis the rpoB-3s primer should be used. For each template to be sequenced, aliquot 3.0ul of each of the nucleotide termination mixes into four separate tubes marked <A>, <C>, <G> and <T> and store on ice until the sequencing mastermix is prepared.

### Cycle sequencing mastermix

rpoB template	2.0ul
10X VGI Sequenace ™ buffer	2.5ul

DMSO 3.5ul
2.5uM dye-sequencing primer 2.0ul
PCR grade water 9.0ul
1:10 diluted Thermosequenase 0.5 ul
total volume 22.0ul

Mix the DMSO and other components in the mastermix well by repeated pipetting (5 times) with a micropipette. Store the mastermix on ice until ready to add to the nucleotide termination mixes.

- 13 -

Add 5.0ul of the mastermix to each of the four marked tubes containing the nucleotide termination mixes.

Add 8.0ul lightweight mineral oil to each of the four marked tubes containing the mastermix and nucleotide termination mixes.

Store on ice until ready to load into the thermocycler.

### Parameters for cycle sequencing

1.	Denaturation	94°C	5 minutes	1X
2.	Denaturation	94°C	30 seconds	
	Annealing	60°C	30 seconds	35X
	Extension	72°C	60 seconds	
3.	Extension	72°C	5 minutes	1X
4.	Hold	6°C		

The temperature change during the cycles of the step 2 is desirably set to ramp at a rate of 1°C/sec.

At the end of the cycle sequencing reaction add 6.0ul of the Stop Loading Dye directly to each of the four tubes to stop the sequencing reaction. The sequencing samples are heated at 95°C for 2 minutes and then placed on ice before loading 2.0ul (from a total volume of 14ul) on the CLIPPER<sup>TM</sup> sequencer. The remainder of the sequencing reaction can be stored at -20°C for subsequent use.

The CLIPPER<sup>TM</sup> sequencer is set-up as described in the *OPENGENE*Automated DNA Sequencing System User Manual. Run parameters for the CLIPPER<sup>TM</sup>
sequencer are 54°C/ 1300volts/ 0.5sec sampling/35min run/50% laser power. The samples loaded included 2 ul each of the forward and reverse sequencing reaction products for the target gene, differentially labeled, for example with CY5.0 and CY5.5 cyanine dye labels. Once the run is completed, the base-called data is analysed by comparison of the test sequence to the rpoB sequence database in GENELIBRARIAN<sup>TM</sup>. This sequence alignment compares the test sequence to the standard control sequence and allows sequence ambiguities to be assessed. Once edited the test sequence can be screened for antibiotic resistance-associated mutations using GENELIBRARIAN<sup>TM</sup>.

Testing for multiple types of antibiotic-resistance mutations can be carried out using a hierarchical assay, as summarized in Fig. 2. At present molecular biological methods for the detection of *M. tuberculosis* are only performed on AFB smear-positive sputum samples. These methods serve as confirmatory tests for the presence of *M. tuberculosis*. In addition to these molecular biological methods, the culture-based procedures for *M. tuberculosis* detection (BacTec liquid culture, agar plate and slant cultures) are performed in parallel. AFB smear-negative sputum samples are processed with only the culture-based detection procedures (Figure 1).

In the present invention both AFB smear-positive and smear-negative sputum samples can be processed using both culture-based and molecular biological methods. A limitation of the AFB stain methodology is it's limit of detection. If a sputum sample has a mycobacterial concentration of less than 5000 bacteria/ul the AFB stain will be negative. In addition to this is the observation that the decontamination procedure used to prepare the sputum sample usually kills 10-20% of the mycobacteria present. This would suggest that two-thirds of the AFB smear-negative samples potentially contain mycobacteria. In practice 10-20% of the AFB smear-negative samples are culture-positive for *M. tuberculosis* (Ontario Public Health Laboratory). This level of mycobacteria is easily detected by molecular biological methods and is therefore incorporated in the present invention.

The hierarchy proposed incorporates tests that specifically detect M. tuberculosis (rpoB), detect mutations in genes associated with resistance to the "first-line" antibiotics used to treat M. tuberculosis infections (rpoB, katG, rpsL/s12, PR, embB, pncA)

and detect other species of mycobacteria (23S) in the absence of M. tuberculosis (Figure 2). Group I analyses are performed before both Group II and Group III. Group I analysis will provide information on the antibiotic resistance status to rifampin (rpoB), isoniazid (katG), streptomycin (rpsL/s12) and azithromycin (23S). In addition the rpoB amplification indicates the presence of M. tuberculosis and in the absence of rpoB amplification the 23S sequence allows identification of most of the clinically relevant mycobacterial species. Group II analysis provides information on antibiotic resistance mutations in the "second-line" antibiotics used to treat M. tuberculosis infections namely, isoniazid (PR), ethambutol (embB), pyrazinamide (pncA) and ciprofloxacin (gyrA). Group III contains gene targets in which mutations associated with antibiotic resistance are infrequently found. This protocol permits specific gene targets to be examined according to the local treatment procedures since the both antibiotics used to treat M. tuberculosis infections, and thus the associated antibiotic resistance mutation patterns, vary geographically. As shown in Figure 2 the culture-based methods are performed in parallel. The molecular biological methods would permit the identification of M. tuberculosis from both AFB smearpositive and smear-negative sputum samples and further provide information on the antibiotic resistance profile of these samples well in advance of current culture-based methods. This information would be crucial to the initiation of appropriate and effective antibiotic treatment regimens for M. tuberculosis infections.

### Examples

A pool of DNA samples from antibiotic-sensitive *M. tuberculosis* isolates was obtained from the LCDC, Health and Welfare Canada. Ottawa, Ontario. Wild-type sequence traces, for all gene targets known to harbor mutations in antibiotic-resistant *M. tuberculosis*, were generated.

A panel of DNA samples from five phenotypic streptomycin-resistant *M.* tuberculosis isolates was obtained from the Public Health Laboratory, Ontario Ministry of Health, Toronto, Ontario. These DNA samples were examined for antibiotic resistance-associated mutations in all 10 antibiotic gene targets listed above. Streptomycin resistance-associated mutations were detected in the rpsL/s12 gene in four isolates. Parallel antibiotic resistance-associated mutations in the rpoB (rifampin), katG (isoniazid), PR (isoniazid),

embB (ethambutol), pncA (pyrazinamide) and gyrA (ciprofloxacin) genes were also identified which underscores the importance of examining all the gene targets for first-line antibiotics used in the treatment of *M. tuberculosis*. A summary of the results is shown in Table 1.

The following references are cited herein and are incorporated herein by reference for all states which allow such incorporation.

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- C Katsukawa et al. (1997). Characterisation of the rpsL and rrs genes of streptomycin-resistant clinical isolates of Mycobacterium tuberculosis in Japan.

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- MA Lety et al. (1997). A single point mutation in the embB gene is responsible for resistance to ethambutol in Mycobacterium smegmatis. Antimicrob Agents Chemother 41: 2629-2633.
- A Scorpio et al. (1997). Characaterisation of pncA mutations in pyrazinamideresistant Mycobacterium tuberculosis. Antimicrob Agents Chemother 41: 540-543.

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OPH#4 bp/codon/aa	tcg553itg, Ser553Leu	wt	N	w	aag88agg, Lys98Arg	w	w	w t	agc95acc, Ser95Thr	*
OPH#3 bp/codon/aa	cac526gac, His526Asp	agc513acc, Ser513Thr	1 %	3	aag43agg, Lys43Arg	w	w	att133aat, lle133Asn	agc95acc, Ser95Thr	w t
OPH#2 bp/codon/aa	tcg553ttg, Ser553Leu	agc513acc, Ser513Thr	w	<del>1</del> 3	aag43agg, Lys43Arg	w t	glc292ttc, val292phe	\$	agc95acc, Ser95Thr	**
OPH#1 bp/codon/aa	cac526tac, His526Tyr	agc513acc, Ser513Thr	g541a	w	wt	w	*	tcc65tct, Ser65Ser	agc95acc, Ser95Thr	* 1
	gene (antiolotic) rpoß (rilampin)	katG.1 (isoniazid)	oxyR-ahpC PR (isoniazid)	fabG (isoniazid)	rpsUs12 (streptomycin)	16s/rrs (streptomycin)	embB (ethambutol)	pncA (pyrazinamide)	gyrA (ciprofloxacin)	23s (azithromycin)